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2 **CO<sub>2</sub>-ENRICHED, LOW, AND VERY LOW, VAPOR PRESSURE**  
3 **LIQUID HYDROCARBON FUELS**  
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6 **TECHNICAL FIELD**  
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8 This invention pertains generally to the physical composition formed by  
9 the combination of carbon dioxide (CO<sub>2</sub>) absorbed in liquid hydrocarbon fuels  
10 having low, and very low vapor pressures. The invention includes fuels used in  
11 fuel-powered engines and devices, such as: diesel and jet aircraft engines, fuel-oil  
12 burning devices such as domestic and commercial heating systems, or electrical  
13 power generation facilities, and resid-oil burning engines such as those used in  
14 large cargo ships.  
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16 **BACKGROUND OF THE INVENTION**  
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18 More particularly, the present invention is a CO<sub>2</sub>-enriched, low, or very low,  
19 vapor pressure hydrocarbon fuel having objectives and advantages that are  
20 important to the fuel industry, to the environment, and to fuel economics. The  
21 present invention:

- 22 a.) provides improved combustion characteristics;  
23 b.) promotes fuel micro-droplet production;  
24 c.) reduces exhaust soot particulate;  
25 d.) reduces exhaust carcinogens;  
26 e.) may be effective in reducing the infrared 'footprint' of the exhaust of  
27 military aircraft and vehicles;  
28 f.) de-oxygenates fuel (including at NTP) during the CO<sub>2</sub>/fuel mixing  
29 process;  
30 g.) provides a means to reduce the complexity and cost of fuel tank inerting  
31 systems;

- 1 h.) provides a cost-effective interim emissions-reducing technology for  
2 existing engines that can readily be implemented, particularly in cities having  
3 high atmospheric soot particulate;
- 4 i.) provides potential tax credit relief to fuel companies for accelerated  
5 ground-based vehicle and aircraft soot particulate reductions;
- 6 j.) provides a means to reduce fuel viscosity (when desired);
- 7 k.) may be effective in cleaning fuel-injection systems;
- 8 l.) may be effective in improving fuel economy;
- 9 m.) which, through improved fuel economy may provide a net reduction in  
10 CO2 production (CO2 in exhaust / per gallon);
- 11 n.) which, through improved fuel economy may provide a cost-savings in  
12 fuel-use which is near, equal to, or greater than, the combined cost of CO2,  
13 CO2-enriched fuel / technology licensing, and systems / equipment used for fuel  
14 tank ullage inerting;
- 15 o.) which can utilize an industrial CO2 that is recovered (re-cycled) from  
16 industrial stacks and vents
- 17 p.) provides new enhanced fuel transferring and storing safety;
- 18 q.) acts as a 'self-inerting fuel';
- 19 r.) uses fuel as a 'weightless container' for transferring and storing  
20 substantial volumes of CO2;
- 21 s.) provides a new means for safely extending Jet-A fuel supplies by mixing  
22 in percentages of JP-4 or naphthas into CO2-enriched Jet-A (e.g. during fuel  
23 shortages, fuel embargoes, or in times of national security if Jet-A supply needs to  
24 be increased, for example, during war, or other international crisis).
- 25 t.) provides a sufficient volume of CO2 in fuel (including at NTP) such that  
26 the CO2 will serve as an inerting medium in the ullages of fuel tanks to which  
27 such fuel is transferred and stored, or where such gas is extracted from fuel  
28 derived from such tanks and is immediately, or is stored and subsequently,  
29 directed back into that fuel tank's ullage;
- 30 u.) is transferrable and storable in, existing fuel distribution systems /  
31 equipment such as those used at airports and other re-fueling terminals;
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1 vacuum state will pull out lighter ends (such as butane) from the fuel, which will  
2 adversely affect fuel performance. Continued exposure to a vacuum will remove  
3 the middle range of hydrocarbon molecules and ultimately the heavier range of  
4 molecules.

5 The Jones patent intent was a vacuum removal of one gas and provide positive  
6 pressure (greater than 1 atmosphere) for placement and maintenance of another  
7 gas (CO<sub>2</sub>), a means to remove oxygen from fuel that might otherwise degas into  
8 the vapor space (ullages) of aircraft fuel tanks containing the Jones'-processed  
9 aviation gasoline.

10 By contrast, the composition of the present invention can be achieved with gas-  
11 fuel mixing at NTP (normal temperature and pressure) i.e. without negative and  
12 positive pressure mixing stages and uses the CO<sub>2</sub> as an inerting medium. It is also  
13 noted that contemporary aviation gasoline now contain anti-oxidants that are  
14 inexpensive and do not require the fuel vacuum and pressurization stages needed  
15 to process the Jones' fuel. Whereas this invention is based upon hydrocarbon fuel  
16 having contemporary needs which it satisfies..

17 CO<sub>2</sub> solubility in contemporary grades of hydrocarbon fuels, show that CO<sub>2</sub>  
18 concentrations in the range Jones specifies would require constant positive-  
19 pressure (if not hi-pressure) storage and handling conditions. Thus, the only way  
20 to maintain Jones' 100%-300% concentrations of CO<sub>2</sub> in modern aviation  
21 gasoline would be to make, transfer, distribute, and store, the fuel continuously  
22 under impractical (or as Jones says "super atmospheric") pressures. It is widely  
23 known that virtually all fuel tanks of diesel trucks, aircraft and other ground-  
24 based vehicles, are vented to allow for expansions and contractions of fuel-tank  
25 ullages caused by altitude and temperature changes, and by fuel usage. Thus, it  
26 would be necessary to retrofit all such vehicles with unvented pressure-capable  
27 tanks in order to store aviation gasoline having the Jones' concentrations of CO<sub>2</sub>  
28 (up to and including the "five atmospheres" he specifies). This goal would be  
29 especially challenging in that a commercial aircraft will typically experience a  
30 change of four atmospheres in the various phases of a single flight (which is why  
31 their tanks must be vented). It would also be necessary to have specially equipped  
32 fuel transportation vehicles for the delivery of Jones' pressurized gasoline.

Such prerequisites to the processing, transferring, transportation, handling, and use of Jones' aviation gasoline again, are evidence that Jones did not achieve a practical physical composition patent, especially as it pertains to contemporary technology. Rather, the Jones' physical composition can only exist with the employment of unconventional physical apparatus that in the contemporary fuel market are not only unnecessary, they would require a costly re-building of entire fuel distribution and storing systems.

By contrast, the present invention provides gas-fuel mixtures with fuels other than gasoline (hydrocarbon fuels with low, and very low, vapor pressures at NTP), where such mixtures are attainable and/or storable at NTP having effective concentrations of CO<sub>2</sub> that provide new fuel safety-enhancing and improved combustion advantages. These advantages are achieved without Jones' hi-pressure transportation, handling and storage conditions.

Lastly, the Jones patent describes a safety-enhancing aspect of his fuel, which, assuming the gas concentrations he specifies were attainable, is intended to reduce the danger of fuel fires in the event of catastrophic fuel tank ruptures. For example, the Jones' fuel is described as being advantageous during aircraft crashes, or during fuel tank ruptures caused by bullets piercing the tanks in times of war. In contrast, the practicably attained physical composition of the present invention is comprised of CO<sub>2</sub>-enrichment of low, or very low, vapor pressure fuels, whose safety-enhancing characteristics are only intended to inert the contents of the ullages of fuel tanks that are intact (i.e. non-ruptured).

#### **SUMMARY OF THE INVENTION**

Vehicles and devices that burn low, and very low, vapor pressure hydrocarbon fuels have been in use for over a century and the chemistry composition of such fuels (including relatively newer jet fuels) have largely remained the same since their commercialization. While such fuels offer a potent source of power to the respective engines or devices that utilize them, they also have drawbacks that the physical composition of the present invention can help to reduce. For instance, it has recently been declared by the California State Air

1 Quality Board that the black soot emissions that emanate from a diesel engine  
2 (used in cars, trucks, heavy equipment, trains, ships and the like), and the soot  
3 from jet turbines, are carcinogenic. Thus, practicably attainable fuel technology,  
4 which improves diesel and jet fuel combustion, particularly during its  
5 acceleration phase (highest emission emitting phase), and otherwise reduces such  
6 emissions, provides an important and timely solution to this environmental and  
7 health-related concern. Recent tests conducted with California State-Approved  
8 Infrared Test Equipment concluded that the improved fuel of the present  
9 invention reduces harmful emissions by as much as 60% during repeatable  
10 accelerations on an unmodified diesel engine when using an optimum and/or  
11 controllable concentration of CO2 absorbed within Diesel #2 fuel. Further testing  
12 using a standardized EPA test produced similar reductions in soot.

13 Alternatively, by controlling concentrations of CO2 absorbed within low  
14 vapor pressure hydrocarbon fuels, the physical composition of the present  
15 invention is also effective in enhancing the safety of the fuel when it is stored in  
16 storage receptacles, whether such receptacles are stationary or reside in any one  
17 or more of a variety of liquid hydrocarbon fuel-powered vehicles. For instance,  
18 hydrocarbon-based fuels can evaporate fuel into a vapor space, or ullage, of the  
19 fuel receptacle in which they reside. These evaporated vapors are usually low  
20 molecular weight hydrocarbons which mix with the air in the ullage, and under  
21 certain conditions have proven to be dangerously explosive. Even JP-4 (a military  
22 aircraft fuel which is mostly kerosene with some low molecular weight naphtha)  
23 usually has a flammable ullage above the fuel and this volatile chemical condition  
24 has had deadly consequences. FAA experts have concluded that the mid-air  
25 explosion of TWA FLT 800 out of New York was due to flammable vapors which  
26 were emitted from relatively heated Jet-A fuel located in the plane's center fuel  
27 tank.

28 Enhanced safety in the storing and/or transporting of these fuels can be  
29 achieved by reducing the amount of air which can enter, or otherwise reside  
30 above the fuel of such fuel tanks by displacing or replacing the air with an inert  
31 gas concentration that will no longer support combustion. For example,  
32 approximately 40% or greater concentration of CO2 will effectively suppress fuel

ullage-combustibility under most operating conditions. The CO<sub>2</sub>-enriched hydrocarbon fuels of the present invention provide this desirable combustion-suppressive condition in two ways. In the first method, the concentration of inert-gas residing in a particular fuel exceeds that fuel's equilibrium CO<sub>2</sub>/fuel state and consequently degasses excess CO<sub>2</sub> from the fuel under known conditions, such as: the passage of time, increases in temperature, agitation of the fuel, and/or a change in relative pressure--such as the ascent of a commercial aircraft to a cruising altitude. In this first method, CO<sub>2</sub>-enriched fuel can be transferred to a fuel tank such that a known excess concentration of CO<sub>2</sub> degasses from the fuel during re-fueling; for example when the fuel tank is only being partially filled with fuel (and the excess degassed CO<sub>2</sub> serves to inert the ullage above the fuel level). In the second method, CO<sub>2</sub>-enriched fuel is pumped from the fuel tank and a gas-scavenging stage of the pump extracts CO<sub>2</sub> from the fuel and the extracted CO<sub>2</sub> is either directed back into the same fuel tank (e.g. its ullage), or is stored in one or more CO<sub>2</sub> storage receptacles for subsequent fuel tank inerting purposes (as described in co-pending patents). In either case, the intended equilibrium state can deliberately be exceeded (without the vacuum stage described in the Jones patent). For example, CO<sub>2</sub> can be mixed in the fuel under controllable positive pressures or under such pressures with agitation (as described in co-pending patents), in which case the fuel is still transferable and storable in conventional fuel delivery systems, and the excess gas will subsequently degas at predictable rates and/or volumes. For example, an aircraft having a shorter duration flight could be fueled with a CO<sub>2</sub>-enriched Jet-A fuel having a gas-to-liquid ratio of CO<sub>2</sub> that is absorbed at a substantially higher ratio, that is achieved by employing a higher mixing pressure of the CO<sub>2</sub> in the liquid fuel, to promote faster degassing than with fuel which is mixed at lower, or ambient, pressures (suitable for longer flights).

It is also noted, that in consideration of the environment, the quantity of CO<sub>2</sub> necessary to inert a commercial aircraft, such as a Boeing 747 flight of 6.5 hours and a distance of 3000 nautical miles, is equal to the amount of CO<sub>2</sub> emitted during just a few seconds of engine exhaust from the flight. Furthermore, the present invention includes the mixing of a commercial grade of CO<sub>2</sub> that has

1 been recovered (re-cycled) from high-CO<sub>2</sub>-content industrial stacks and vents.  
2 Moreover, standardized EPA testing, exhaust pyrometer testing, and engine RPM  
3 analysis have each indicated that improved fuel performance may be caused by  
4 the CO<sub>2</sub>-enriched fuels of the present invention (the improved fuel economy  
5 being caused by the CO<sub>2</sub> reducing fuel droplet size, and/or by a cleaning of fuel  
6 injection components). For a frame of reference: each one half percent  
7 improvement in fuel economy would reduce the 747's 3000 mile flight output of  
8 CO<sub>2</sub> by 2200 pounds (this is several times the amount of CO<sub>2</sub> needed to inert the  
9 fuel tanks during the entire flight, and thus, would represent a net reduction in  
10 CO<sub>2</sub> production).

11 Thus practicably attained CO<sub>2</sub>-enriched liquid hydrocarbon fuels are  
12 provided which overcome significant shortcomings of gas-fuel mixtures requiring  
13 special processing, handling, and unconventional physical apparatus, and which  
14 achieve an enhanced safety fuel, as well as an improved combustion fuel, and do  
15 so using inexpensive (and optionally recycled) CO<sub>2</sub>.

#### 16 17 **BRIEF DESCRIPTION OF THE DRAWINGS**

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19 FIGURE 1 shows a graph of the volume of "Carbon Dioxide" (CO<sub>2</sub>) that  
20 can be mixed, or absorbed, in a volume of low vapor pressure liquid hydrocarbon  
21 fuels, namely "JP-4 / Jet-B", "JP-8 / Jet-A", "JP-7", and "JP-5".

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23 FIGURE 2 is a diagrammatical depiction of a barrel of crude oil  
24 illustrating the hydrocarbon fuel constituents of the barrel, and ranges of fuel in  
25 the barrel representing "Low, and Very Low, Vapor Pressure Fuels", "High Vapor  
26 Pressure Fuels", "Jet-A1 or JP-8", "JP-3", "JP-4", and "JP-5".

#### 27 28 **DETAILED DESCRIPTION OF THE INVENTION**

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30 The graph in FIG. 1 plots volume ratio and depicts a range of the volume  
31 of CO<sub>2</sub> that can be mixed, or absorbed, in a volume of low vapor pressure liquid  
32 hydrocarbon fuels, namely "Diesel, JP-4 / Jet-B", "JP-8 / Jet-A", "JP-7", and "JP-



1 5 and shows that as temperature goes up the volume ratio of the CO<sub>2</sub> within the  
2 fuels goes down. For example, at -20°C about 1.7 volumes of CO<sub>2</sub> can be  
3 absorbed in one volume of JP1- Jet-A. At just under 50°C an approximate 1:1  
4 ratio of gas to fuel is possible. Commercial vehicle and aircraft fuels such as  
5 diesel#2 and Jet-A have a similar composition and consequently similar  
6 absorption ratios of gas to fuel are possible.

7 Several physical factors are known to change the ratio of inert-gas which  
8 is mixable in, and/or retainable within, a particular type of low, or very low, vapor  
9 pressure liquid hydrocarbon fuel; the CO<sub>2</sub> gas is physically absorbed without  
10 reacting chemically with the fuel. These factors can be controlled, or predicted,  
11 and in numerous instances require no additional physical structure, or equipment,  
12 in order to occur. These factors include: temperature change, pressure differential,  
13 agitation, time, and, convection of the mixed or mixing fuel. Thus, a factor such  
14 as a known temperature change during a mixing phase, or during a degassing  
15 phase, will have predictable results which can be advantageously used. For  
16 example, cooler CO<sub>2</sub>-enriched liquid hydrocarbon fuel may be stored in relatively  
17 cooler underground tanks, with the knowledge that as the fuel becomes warmer it  
18 will degas faster at a known rate. Similarly, agitation during a mixing phase or  
19 during a degassing phase will have predictable results. For instance, after  
20 refueling a vehicle or aircraft will encounter fuel agitation as it rolls over the  
21 natural series of bumps such as one encounters on roads (car), or during a 40  
22 second or so takeoff roll (commercial aircraft). Alternatively, a pressure  
23 differential can be exploited to accelerate gas absorption in, and/or desorption  
24 from, fuels, as mentioned previously in the case of aircrafts having different flight  
25 durations. Time is also a known factor that can be used advantageously to predict  
26 absorption or desorption rates, including rates when also affected by any of the  
27 above-mentioned factors that can change the ratio of the physical composition. Or  
28 CO<sub>2</sub>-enriched fuel can be made with a gas/fuel ratio that remains fairly stable  
29 through all phases of a flight (i.e. through all changes in altitude).

30 In the case of the physical composition being used as a vapor combustion-  
31 suppression fuel, it can be beneficial to mix CO<sub>2</sub> in fuels such as diesel #2 or Jet-  
32 A in a ratio that exceeds the fuel's equilibrium state of the gas/fuel mixture (such

as that exceeding the 50% ratio previously mentioned). For example, a 1.5:1 gas/fuel ratio might be employed where 0.75 of a fuel's CO<sub>2</sub> volume is meant to degas from the fuel under relatively ambient conditions, and more CO<sub>2</sub> can degas, or be extracted from the fuel, according to one or more of the factors or methods previously mentioned. Since an ullage in a tank containing a typical hydrocarbon-based fuel can be inerted by containing approximately 35% or more volume of CO<sub>2</sub> therein, it can be seen that a sufficient volume of inert-gas can be retained within the fuel to in effect act as a "self-inerting-fuel".

In the case of the application of these physical compositions providing an improved combustion and/or emission-reducing fuel, it can be beneficial to mix the CO<sub>2</sub> with the hydrocarbon fuel in less than 1:1 to 3:1 gas/liquid ratio of CO<sub>2</sub> to fuel (e.g. a range from 0.1:1 to 1:1) gas/fuel ratios in the approximate range of 15-25% that have been shown to reduce harmful particular emissions in diesel #2 fuel by as much as 60%. The various CO<sub>2</sub>/fuel ratios may indicate that minute inert-gas bubbles can form in 'micro-droplets', and upon reaching the surface of the droplet can disperse the droplets into smaller ones which creates a larger mixing surface for fuel and air molecules for optimized dispersion of the fuel which results in improved (and more complete) combustion in an engine or fuel-burning device. This benefit may be of further advantage during cold weather, or during cold-engine starting.

Additionally, liquid fuel convection is a means of replacing surface fuel molecules so that they may absorb CO<sub>2</sub> molecules from a blanket of CO<sub>2</sub> gas above the liquid surface. Thus, the factors of convection and time, or convection and fuel temperature and/or CO<sub>2</sub> temperature, or other factors mentioned above can be predictable and used advantageously to attain desired CO<sub>2</sub>/fuel concentrations and results. Without the surface layer of CO<sub>2</sub> the same methods create degassing of the CO<sub>2</sub>.

Because a CO<sub>2</sub>-enriched low vapor pressure fuel can be easily and practicably attained by a number of controllable methods, with the outcome being a CO<sub>2</sub>/fuel mixture that is transferrable and storable in existing fuel distribution systems, the present invention provides a fuel that can be easily made at a number of junctures in the fuel distribution system. For instance, at an airport fuel farm, at

any one or more of a variety of re-fueling terminals, in underground storage tanks, on fuel tankers, fuel barges, on a ship, and the like.

Figure 2 shows a diagrammatical depiction of a barrel of crude oil illustrating the hydrocarbon fuel constituents of the barrel. Ranges of fuel in the barrel representing low vapor pressure fuels are bracketed, and those representing high vapor pressure fuels are separately bracketed. It can also be seen in Fig. 2 that Jet-A1 or JP-8 can be derived from 10% of the crude, JP-3 from 50% of the crude, JP-4 from 25% of the crude, and JP-5 from 2% of the crude. An important benefit of the present invention pertains to fuel supply, particularly during a national emergency such as a fuel shortage caused by any one of a number of factors, such as an international crisis, a war, a fuel embargo, or an international trust type of control over the supply / cost of fuels. In any such instance, for example a shortness of supply in Jet-A fuel could threaten national security (since this fuel is used in military aircraft). Heretofore, JP-4 fuel has been eliminated from fuel supplies, due to proven fuel tank ullage volatility associated with its higher vapor pressures. The range of CO<sub>2</sub>-enriched fuels of the present invention, and the range of CO<sub>2</sub> that can be absorbed within low, and very low vapor pressure fuels, make it possible to significantly increase jet fuel supply in times of emergency by providing the means to extend CO<sub>2</sub>-enriched Jet-A supplies by mixing in percentages of JP-4 or Naphthas (and increasing CO<sub>2</sub> as needed), or increasing other usable jet fuel supplies by mixing suitable concentrations of CO<sub>2</sub> in one or more non-Jet-A "JP" fuels. Such an approach can alternatively be employed in a non-crisis situation, for instance where a government / country may be close to being self-sufficient in one or more fuel and could attain self-sufficiency by a prudent mixing of jet fuels and CO<sub>2</sub>.

The fact that carbon particulate is reduced in the engine exhaust means that any infra-red emissions from this source are also minimized. This can have value to military exhaust cloaking efforts.

Although the present invention has been described with a certain degree of particularity, it is understood that the present disclosure has been made by way of

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example, and changes in detail or structure may be made without departing from the spirit of the invention as defined in the appended claims.